

This Page Is Inserted by IFW Operations  
and is not a part of the Official Record

## **BEST AVAILABLE IMAGES**

Defective images within this document are accurate representations of the original documents submitted by the applicant.

Defects in the images may include (but are not limited to):

- BLACK BORDERS
- TEXT CUT OFF AT TOP, BOTTOM OR SIDES
- FADED TEXT
- ILLEGIBLE TEXT
- SKEWED/SLANTED IMAGES
- COLORED PHOTOS
- BLACK OR VERY BLACK AND WHITE DARK PHOTOS
- GRAY SCALE DOCUMENTS

**IMAGES ARE BEST AVAILABLE COPY.**

**As rescanning documents *will not* correct images,  
please do not report the images to the  
Image Problem Mailbox.**



GenCore version 4.5  
Copyright (c) 1993 - 2000 - CompuGen Ltd.

OM nucleic - nucleic search, using sw model

Run on: April 19, 2002, 19:19:31 ; Search time 344.5 Seconds

(without alignments)  
10434.731 Million cell updates/sec

Title: US-09-731-457B-3

Perfect score: 4193  
Sequence: 1 gtgagcttcgtcgcgcgtcgtt.....agtttaccataaagtag 4193

Scoring table: OLIGO\_MNC  
Gapop 60.0, Gapext 60.0

Searched: 930621 seqs, 428662619 residues

Word size: 0

Total number of hits satisfying chosen parameters: 989696

Minimum DB seq length: 0

Maximum DB seq length: 50

Post-processing: Listing first 45 summaries

Database:

N.Geneseq\_1101.\*  
1: /SID2/gcgdata/geneseq/geneseqn/NA1980.DAT:\*  
2: /SID2/gcgdata/geneseq/geneseqn/NA1981.DAT:\*  
3: /SID2/gcgdata/geneseq/geneseqn/NA1982.DAT:\*  
4: /SID2/gcgdata/geneseq/geneseqn/NA1983.DAT:\*  
5: /SID2/gcgdata/geneseq/geneseqn/NA1984.DAT:\*  
6: /SID2/gcgdata/geneseq/geneseqn/NA1985.DAT:\*  
7: /SID2/gcgdata/geneseq/geneseqn/NA1986.DAT:\*  
8: /SID2/gcgdata/geneseq/geneseqn/NA1987.DAT:\*  
9: /SID2/gcgdata/geneseq/geneseqn/NA1988.DAT:\*  
10: /SID2/gcgdata/geneseq/geneseqn/NA1989.DAT:\*  
11: /SID2/gcgdata/geneseq/geneseqn/NA1990.DAT:\*  
12: /SID2/gcgdata/geneseq/geneseqn/NA1991.DAT:\*  
13: /SID2/gcgdata/geneseq/geneseqn/NA1992.DAT:\*  
14: /SID2/gcgdata/geneseq/geneseqn/NA1993.DAT:\*  
15: /SID2/gcgdata/geneseq/geneseqn/NA1994.DAT:\*  
16: /SID2/gcgdata/geneseq/geneseqn/NA1995.DAT:\*  
17: /SID2/gcgdata/geneseq/geneseqn/NA1996.DAT:\*  
18: /SID2/gcgdata/geneseq/geneseqn/NA1997.DAT:\*  
19: /SID2/gcgdata/geneseq/geneseqn/NA1998.DAT:\*  
20: /SID2/gcgdata/geneseq/geneseqn/NA1999.DAT:\*  
21: /SID2/gcgdata/geneseq/geneseqn/NA2000.DAT:\*  
22: /SID2/gcgdata/geneseq/geneseqn/NA2001.DAT:\*

Pred. No. is the number of results predicted by chance to have a score greater than or equal to the score of the total being printed, and is derived by analysis of the total score distribution.

#### SUMMARIES

Result No.	Score	Query Match	Length DB	ID	Description
1	28	0.7	39	18	AAAT65714
2	28	0.7	48	21	AAAT7335
3	28	0.7	50	13	AAQ34003
4	27	0.6	45	18	AAAT65737
5	26	0.6	29	13	AAQ3630
6	25	0.6	35	13	AAQ3713
7	25	0.6	44	21	AAAT7337
8	24	0.5	40	18	AAAT65736
9	23	0.5	50	18	AAAT65741
10	22	0.5	27	13	AAQ34077
11	21	0.5	24	22	AAH39357

C	12	21	0.5	40	18	AAAT65743
C	13	20	0.5	20	16	AAQ82617
C	14	20	0.5	20	16	AAQ82618
C	15	20	0.5	21	19	AAAT58080
C	16	20	0.5	21	19	AAAT58080
C	17	19	0.5	27	21	AAAT65736
C	18	19	0.5	35	13	AAQ33695
C	19	19	0.5	37	13	AAQ33695
C	20	19	0.5	37	13	AAQ33695
C	21	19	0.5	46	18	AAAT65709
C	22	18	0.4	50	13	AAQ34167
C	23	18	0.4	24	14	AAQ40668
C	24	18	0.4	32	13	AAQ34119
C	25	18	0.4	37	13	AAQ33698
C	26	18	0.4	43	18	AAAT65788
C	27	18	0.4	44	21	AAAT65788
C	28	17	0.4	23	21	AAAT65788
C	29	17	0.4	25	21	AAAT65788
C	30	17	0.4	40	13	AAQ34094
C	31	17	0.4	40	21	AAAT65797
C	32	17	0.4	42	18	AAAT65797
C	33	17	0.4	47	21	AAAT65797
C	34	17	0.4	49	13	AAQ34122
C	35	16	0.4	19	17	AAAT30412
C	36	16	0.4	19	17	AAAT30412
C	37	16	0.4	20	14	AAQ48546
C	38	16	0.4	20	17	AAAT30427
C	39	16	0.4	20	22	AAAT65797
C	40	16	0.4	23	22	AAAT65797
C	41	16	0.4	24	21	AAAT65797
C	42	16	0.4	26	13	AAQ34131
C	43	16	0.4	27	22	AAAT65797
C	44	16	0.4	27	22	AAAT65797
C	45	16	0.4	34	13	AAQ33761

#### ALIGNMENTS

RESULT 1	AAAT65714/c	AAAT65714 standard; DNA; 39 BP.
XX	AAAT65714/c	
AC	AAAT65714/c	
XX	AAAT65714/c	
DT	17-JUN-1997 (first entry)	
XX	Repeat sequence from polymorphic marker clone Mfd12.	
DE	Polymorphism; repeat sequence; genetic marker; primer; amplification;	
XX	PCR; polymerase chain reaction; paternity; maternity; human; pedigree;	
KW	linkage analysis; genetic disease; animal; plant; breeding; locus;	
KW	hybridisation; chromosome; ds.	
XX		
OS	Homo sapiens.	
XX		
PN	US5582979-A.	
XX		
PD	10-DEC-1996.	
XX		
PF	21-APR-1989; 89US-0341562.	
XX		
PR	05-SEP-1991; 91US-0754351.	
PR	21-APR-1989; 89US-0341562.	
PR	04-APR-1994; 94US-0222177.	
XX		
PA	(MARS-), MARSHFIELD CLINIC.	
XX		
PI	Weber JL;	
XX		
DR	WPI; 1997-042299/04.	
XX		
PT	Detection of polymorphic genetic markers of the form	

Repeat sequence fr  
Chromosome 11 (loc  
Chromosome 11 (loc  
ICAM-1 antisense o  
Human ICAM-1, E-se  
Human ASTH11 5' re  
Microsatellite seq  
Microsatellite seq  
Repeat sequence fr  
Sequence of a micr  
LPA-12 in vitro mu  
H. discus derived  
Sequence of a micr  
Microsatellite seq  
Repeat sequence fr  
H. discus derived  
Human ASTH11 5' re  
Mahogany protein g  
Sequence of a micr  
H. discus derived  
Repeat sequence fr  
Human map-related  
Sequence of a micr  
Compound simple se  
Primer TEL:114019  
HPV E6/7 region pr  
Compound simple se  
Human MEXK2 antisense  
Human FKL1 gene tr  
H. discus derived  
Sequence of a micr  
Microsatellite seq  
Soybean 240017 reg  
Microsatellite seq









DR WPI; 1992-284684/34.

PT Polymorphic bovine DNA markers - used in genetic identification,  
PF gene mapping, and selective breeding

XX  
XX PS Table 7; Page 362; 517pp; English.

CC The sequence is that of a bovine microsatellite sequence obt. by  
CC screening a library of bovine MboI DNA fragments of between  
CC 250 and 500 bp with an (AC)<sub>15</sub> and a (TC)<sub>15</sub> oligonucleotide probe.  
CC One out of 30 clones cross-hybridised. Assuming independent  
CC distribution of microsatellites and MboI sites, the frequency of  
CC (TC)<sub>n</sub> >9 microsatellites in the bovine genome is estimated at >100,  
CC 000. The sequence information for ca. 230 such bovine microsatellites  
CC is summarised in the specification and indexed herein (see below).  
CC The sequences upstream and downstream of the microsatellite sequence  
CC were used to generate the required PCR primers for *in vitro*  
CC amplification of the corresp. microsatellite (using the program  
CC OPIPprim). The microsatellites may be used to identify individuals,  
CC for percentage testing, and in the genetic mapping of economic trait  
CC loci or genes involved in the determination of economically important  
CC traits esp. in cattle, to allow selective breeding.  
CC See also AAQ33501-34437.

SX Sequence 27 BP; 1 A; 0 C; 13 G; 13 T; 0 other;

QY Query Match 0.58; Score 22; DB 13; Length 27;  
Best Local Similarity 100.0%; Pred. No. 3.3;  
Matches 22; Conservative 0; Mismatches 0; Indels 0; Gaps 0;

OY 3855-gtgcgtgcatgtgtgtgtgtgt 3876  
|||||  
DB 1 gtgcgtgcatgtgtgtgtgtgt 22

RESULT 11-

AAH39357  
ID AAH39357 standard; DNA; 24 BP.

XX  
XX AC AAH39357;

XX  
XX DT 14-AUG-2001 (first entry)

DE SNP specific upper PCR primer SEQ ID 2153.

XX  
XX Single nucleotide polymorphism; SNP; single nucleotide primer extension;  
KM SNEB; genotyping; agammaglobulinemia; diabetes insipidus; cancer;  
KM Leach-Nyman syndrome; muscular dystrophy; familial hypercholesterolaemia;  
KM polycystic kidney disease; osteogenesis imperfecta; autoimmune disease;  
KM acute interstitial porphyria; rheumatoid arthritis; multiple sclerosis;  
KM inflammation; forensic investigation; paternity analysis; PCR primer; ss.

OS Homo sapiens.

XX  
XX PN WC200129262-A2.

XX  
XX PD 26-APR-2001.

Pf 13-OCT-2000; 2000WO-US28436.

XX PR 15-OCT-1999; 99US-0160096.

XX PA (ORCH-) ORCHID BIOSCIENCES INC.

XX PI Picoult-Newburg L., Pohl M;

XX WPI; 2001-230930/30.

XX  
XX New genotyping oligonucleotide, useful for detecting the presence,  
PT absence or identity of single polynucleotide polymorphism in a nucleic  
acid sample -

PS Claim 1; Page 60; 83bp; English.

XX Sequences AA37205 - AA40944 represent PCR primers, single nucleotide  
CC primer extension (SNPE) primers, and the sequences of regions flanking  
CC sites of single nucleotide polymorphisms SNPs. The present invention  
CC includes kits for determining the presence or absence of a SNP, using the  
CC oligonucleotides of the invention. The PCR primers are used to amplify a  
CC SNP flanking sequence, the SNPE primer is used as a genotyping primer.  
CC The oligonucleotides are useful for genotyping a nucleic acid sample by  
CC performing a single-nucleotide primer extension reaction. The  
CC oligonucleotides are useful for determining the presence, absence or  
CC identity of a SNP and for genotyping nucleic acid samples, for e.g. to  
CC assess by association analysis the genotype of an individual or group of  
CC individuals, having a pathological phenotypic trait suspected of being  
CC caused by one or more SNPs. Phenotypic traits include diseases e.g.  
CC agammaglobulinemia, diabetes insipidus, Leisch-Nyhan syndrome, muscular  
CC dystrophy, familial hypercholesterolaemia, polycystic kidney disease,  
CC osteogenesis imperfecta and acute intermittent porphyria. Phenotypic  
CC traits also include symptoms of or susceptibility to multifactorial  
CC diseases of which a component is or may be genetic such as autoimmune  
CC diseases, including, rheumatoid arthritis, multiple sclerosis,  
CC inflammation, cancer, nervous system diseases and infection by pathogenic  
CC microorganism. The method is also useful in forensic investigations and  
CC paternity analysis. The present sequence represents a PCR primer specific  
CC for a human SNP containing DNA sequence.

SQ Sequence 24 BP; 1 A; 0 C; 11 G; 12 T; 0 other;

Query Match 0.5%; Score 21; DB 22; Length 24;  
Best Local Similarity 100.0%; Pred. No. 9.8;  
Matches 21; Conservative 0; Mismatches 0; Indels 0; Gaps 0;

OY 3854 tctgtgtatctgtgtgtgt 3874  
Db 1 tgtgtgtatctgtgtgtgt 21  
|||||

RESULT 12  
AAT65743/C  
ID AAT65743 standard; DNA; 40 BP.  
XX AAT65743;  
AC  
XX 17-JUN-1997 (first entry).  
DT  
XX  
DE Repeat sequence from polymorphic marker clone Mfd42.  
XX  
KW Polymorphism; repeat sequence; genetic marker; primer; amplification;  
KW PCR; polymerase chain reaction; paternity; maternity; human; pedigree;  
KW linkage analysis; genetic disease; animal; plant; breeding; locus;  
KW hybridisation; chromosome; ds.  
XX  
OS Homo sapiens.  
XX  
XX US5582979-A.  
PN  
XX  
PD 10-DEC-1996.  
XX  
PF 21-APR-1989; 89US-0341562.  
XX  
PR 05-SEP-1991; 91US-0754351.  
PR 21-APR-1989; 89US-0341562.  
PR 04-APR-1994; 94US-0222177.  
XX  
PA (MARS-) MARSHFIELD CLINIC.  
XX  
XX  
PI Weber JL;  
XX  
XX WPI; 1997-042299/04.  
DT  
XX  
XX Detection of polymorphic genetic markers of the form  
(C-C-dh)n(dg-dfn) - using novel nucleic acid mols. as primers



```

PS   Disclosure; Column 9-10; 186pp; English.
XX
CC   The invention relates to the isolation of polymorphic repeat sequences
CC   having the sequence (dc-da)n.(dg-dt)n which can be used as genetic
CC   markers. Primers based on these sequences can be used to detect these
CC   repeats, especially for use in e.g. paternity or maternity testing,
CC   human genetic analysis, such as linkage analysis of genetic disease,
CC   commercial animal or plant breeding or pedigree analysis. Clones
CC   containing the repeat sequences were isolated by hybridisation of
CC   chromosome-specific phage libraries with a synthetic poly(dc-da).(dg-dt)
CC   probe. Over 100 repeat blocks were isolated. The inserts from the
CC   clones were amplified by primers AAT65798-766047. Those clones where the
CC   repeat sequence has been determined are shown in AAT65704-797. This
CC   repeat sequence is from the marker clone W6f42 which contains the repeat
CC   sequence having the formula: (CA)167TAC)3.5.
XX
SQ   Sequence 40 BP; 20 A; 19 C; 0 G; 1 T; 0 other;
XX
Query Match          0.5%; Score 21; DB 18; Length 40;
Best local Similarity 100.0%; Pred. No. 9.7;
Matches 21; Conservative 0; Mismatches 0; Indels 0; Gaps 0
OY 3856 tctgtctatgtctgtctgtct 3876
    |||||
    40 TGTGTGTATGTGTGTGTGTGT 20
XX
RESULT 13
AA082617/c
ID   AA082617 standard; DNA; 20 BP.
XX
AC   AA082617;
XX
DE   14-SEP-1995 (first entry)
XX
DT   Chromosome 11 (locus D11S952E) STS primer 339.
XX
KM   sequence sampled mapping; genomic analysis; complex genome mapping;
XX   cosmid library; chromosome 11; sequence tagged site; STS analysis; ss.
XX
OS   Synthetic.
XX
PN   W09429486-A.
XX
PD   22-DEC-1994.
XX
PF   15-JUN-1994; 94WO-US06810.
XX
PR   15-JUN-1993; 93US-0078471.
XX   07-SEP-1993; 93US-0117952.
XX
PA   (SALK ) SALK INST BIOLOGICAL STUDIES.
XX
PI   Evans GA, Smith MW;
XX
WL   1995-036508/05.
XX
PT   Sequencing complex genomes, present as fragments in a cosmid
PT   library - by sequencing end-specific nucleotides of each clone
PT   then correlating with spatial relationship of cosmid, esp. for
PT   mammalian chromosomes.
XX
PS   Example 4; Page 90; 128pp; English.
XX
CC   Sequences were determined from the ends of chromosome 11-specific
CC   cosmids by automated sequencing without intermediate subcloning.
CC   A sample of 371 DNA sequence fragments were determined and of
CC   these, 277 were suitable for STS primer prediction by computer
CC   analysis (using the "Primer" program available from E.Lander, MIT).
CC   The STS and cosmids were mapped by in situ hybridisation, somatic
CC   cell hybrid analysis or both. Using this method, 370 STS specific

```

CC for human chromosome 11 were generated and most of them were  
 CC regionally mapped. This procedure illustrates a novel method for  
 CC sequencing complex genomes, designated "sequence sampled mapping".  
 CC The sequence sampled mapping method is useful for the completion of  
 CC high density sequence-based maps, and ultimately, for the complete  
 CC sequencing of genomic DNA directly from cosmid clones.  
 CC See AA082001-082706 and AA091325-091358 for STS primers.  
 XX  
 S0 Sequence 20 BP; 5 A; 3 C; 6 G; 6 T; 0 other:  
 Query Match 0.5%; Score 20; DB 16; Length 20;  
 Best Local Similarity 100.0%; Pred. No. 29;  
 Matches 20; Conservative 0; Mismatches 0; Indels 0; Gaps 0.  
 CY 4119 caaactgcctcttcgaaa 4138  
 Db 20 CAAACTGCCTCTTCGAAA 1  
 |||||  
 RESULT 14  
 ID AA082618  
 AC AA082618 standard; DNA: 20 BP.  
 XX  
 DT 14-SEP-1995 (first entry)  
 XX  
 DE Chromosome 11 (locus D11S952E) STS-primer 340.  
 KM sequence sampled mapping; genomic analysts; complex genome mapping;  
 KM cosmid library; chromosome 11; sequence tagged site; STS analysts; ss.  
 XX  
 OS Synthetic.  
 XX  
 PM W09429486-A.  
 PD 22-DEC-1994.  
 XX  
 PF 15-JUN-1994; 94WO-US06810.  
 XX  
 PR 15-JUN-1993; 93US-0078471.  
 PR 07-SEP-1993; 93US-0117952.  
 PA (SALK ) SALK INST BIOLOGICAL STUDIES.  
 PI Evans GA, Smith MW;  
 XX  
 DR WPI; 1995-036508/05.  
 XX  
 PT Sequencing complex genomes, present as fragments in a cosmid  
 PT library - by sequencing end-specific nucleotides of each clone  
 PT then correlating with spatial relationship of cosmid, esp. for  
 PT mammalian chromosomes.  
 XX  
 PS  
 XX  
 Example 4; Page 90; 128bp; English.  
 CC Sequences were determined from the ends of chromosome 11-specific  
 CC cosmid by automated sequencing without intermediate subcloning.  
 CC A sample of 371 DNA sequence fragments were determined and of  
 CC these, 277 were suitable for STS primer prediction by computer  
 CC analysis (using the "Primer" program available from E.Lander, MIT).  
 CC The STSs and cosmid were mapped by in situ hybridisation, somatic  
 CC cell hybrid analysis or both. Using this method, 370 STSs specific  
 CC for human chromosome 11 were generated and most of them were  
 CC regionally mapped. This procedure illustrates a novel method for  
 CC sequencing complex genomes, designated "sequence sampled mapping".  
 CC The sequence sampled mapping method is useful for the completion of  
 CC high density sequence-based maps, and ultimately, for the complete  
 CC sequencing of genomic DNA directly from cosmid clones.  
 CC See AA082001-082706 and AA091325-091358 for STS primers.  
 XX  
 S0 Sequence 20 BP; 5 A; 6 C; 5 G; 4 T; 0 other:

Query Match 0.5%; Score 20; DB 16; Length 20;  
 Best Local Similarity 100.0%; Pred. No. 29;  
 Matches 20; Conservative 0; Mismatches 0; Indels 0; Gaps 0;

OY 4024 atcagcctagagcctgact 4043  
 |||||  
 Db 1 atcagcctagagcctgact 20

RESULT 15  
 AAT58080/c  
 ID AAT58080 standard; DNA: 21 BP.

XX AAT58080;  
 XX  
 XX 18-MAR-1997 (first entry)  
 XX  
 XX ICAM-1 antisense oligonucleotide #10.

XX Antisense; pre-mRNA; mature mRNA; vascular defect; tissue defect;  
 KW human intercellular adhesion molecule-1; ICAM-1; inflammation;  
 KW adult respiratory distress syndrome; multiple organ failure; GMI594;  
 KW septic shock; ss.

XX Synthetic.

XX USS580969-A.

XX 03-DEC-1996.

XX 24-JUL-1992; 92US-0918259.

XX 12-OCT-1993; 93US-0136118.

XX 24-JUL-1992; 92US-0918259.

XX (USNA) US SEC OF NAVY.

XX Bradley MO, Hoke GD, Lee C, Williams TJ;

XX WPI; 1997-033603/03.

XX Anti-sense oligo:nucleotide(s) for blocking ICAM-1 mRNA translation  
 PT for treating septic shock, adult respiratory distress syndrome  
 PT etc.

XX Claim 1; Column 21; 16pp; English.

XX The sequences given in AAT58071-85 represent oligonucleotides which are  
 CC antisense to sequences contained in the pre-mRNA or mature mRNA  
 CC transcript of human intercellular adhesion molecule-1 (ICAM-1).  
 CC These oligonucleotides may be used for treating septic shock and the  
 CC manifestations of septic shock, e.g. inflammation, and vascular and  
 CC tissue defects. They are also useful in the treatment of septic  
 CC shock associated diseases, e.g. adult respiratory distress syndrome,  
 CC multiple organ failure etc.

XX Sequence 21 BP; 11 A; 9 C; 0 G; 1 T; 0 other;

Query Match 0.5%; Score 20; DB 18; Length 21;  
 Best Local Similarity 100.0%; Pred. No. 29;  
 Matches 20; Conservative 0; Mismatches 0; Indels 0; Gaps 0;

OY 3849 gctgtgtgtgtgtgtgt 3868  
 |||||  
 Db 20 gctgtgtgtgtgtgtgt 1

RESULT 16  
 AAV38616/c  
 ID AAV38616 standard; DNA: 21 BP.

XX AAV38616;

XX 13-OCT-1998 (first entry)

XX Human ICAM-1, E-selectin, VCAM-1 antisense oligonucleotide.

XX ICAM-1; intracellular adhesion molecule-1; E-selectin; VCAM-1;  
 KW vascular cell adhesion molecule-1; antisense; inflammatory;  
 KW disease; treatment; septic shock; psoriasis; wounds; burns; acne;  
 KW arthritis; organ rejection; inhibition; expression; ss.

XX Synthetic.  
 OS Homo sapiens.

XX WO9824797-A1.

XX 11-JUN-1998.

XX 02-DEC-1996; 96WO-US19194.

XX 02-DEC-1996; 96WO-US19194.

XX (DYAD-) DYAD PHARM CORP.

XX Bradley MO, Hoke GD, Lee C, Williams TJ;

XX WPI; 1998-333253/29.

XX Antisense oligonucleotides to ICAM-1, E-selectin or VCAM-1 - useful  
 PT for treating diseases having an inflammatory component, e.g.  
 PT psoriasis, wounds and septic shock

XX Claim 8; Page 40; 48pp; English.

XX The sequence is that of an antisense oligonucleotide which is  
 CC substantially complementary to at least a portion of the pre-  
 CC or mature RNA transcript of human intercellular adhesion molecule  
 CC (ICAM), E-selectin or vascular cell adhesion molecule (VCAM).  
 CC It can be used to inhibit expression of these proteins. Inhibition  
 CC of these proteins forms the basis for treatment of conditions and  
 CC diseases that have an inflammatory component, e.g. acne, psoriasis,  
 CC arthritis, organ rejection, wounds, burns, septic shock or  
 CC inflammatory complications of septic shock.

XX Sequence 21 BP; 11 A; 9 C; 0 G; 1 T; 0 other;

Query Match 0.5%; Score 20; DB 19; Length 21;  
 Best Local Similarity 100.0%; Pred. No. 29;  
 Matches 20; Conservative 0; Mismatches 0; Indels 0; Gaps 0;

OY 3849 gctgtgtgtgtgtgtgt 3868  
 |||||  
 Db 20 gctgtgtgtgtgtgtgt 1

RESULT 17  
 AAA80358  
 ID AAA80358 standard; DNA: 27 BP.

XX AAA80358;

XX 22-NOV-2000 (first entry)

XX Human ASTH1 5' region polymorphic site, SEQ ID NO:103 (b).

XX ASTH1 locus; ASTH1; ASTH1; human; chromosome 11p; asthma;  
 KW bronchial hyperreactivity; ets family; transcription factor;  
 KW splice variant; genetic predisposition; polymorphism; antibody;  
 KW drug screening; prophylaxis; therapy; diagnosis;  
 KW single nucleotide polymorphism; SNP; ss.

RESULT 18  
AA033695 standard; DNA; 35 BP.  
XX AA033695;  
DT 02-FEB-1993 (first entry)  
DE Microsatellite sequence from clone TGLA127.  
KM PCR selection; primers: OPTIPRIM; breeding; cattle; parentage;  
KW genetic mapping; traits; amplification; ss.  
OS Bos taurus.  
PN MO9213102-A.  
PD 06-AUG-1992.  
PE 15-JAN-1992; 92MO-US00340.  
PF 15-JAN-1991; 91US-0642342.  
PR (GENM-) GENMARK.  
PA Georges M, Massey JM;  
PI WPI; 1992-284684/34.  
DR Polymorphic bovine-DNA markers - used in genetic identification,  
PT gene mapping, and selective breeding  
PS Table 7; Page 208; 517pp; English.

XX The sequence is that of a bovine microsatellite sequence obtd. by  
CC screening a library of bovine MboI DNA fragments of between  
CC 250 and 500 bp with an (AC)15 and a (TC)15 oligonucleotide probe.  
CC One out of 50 clones cross-hybridised. Assuming independent  
CC distribution of microsatellites and MboI sites, the frequency of  
CC (re)n >9 microsatellites in the bovine genome is estimated at >100,  
000. The sequence information for ca. 230 such bovine microsatellites  
CC is summarised in the specification and indexed herein (see below).  
CC The sequences upstream and downstream of the microsatellite sequence  
CC were used to generate the required PCR primers for in vitro  
CC amplification of the corresp. microsatellite (using the program  
CC OPTIPRIM). The microsatellites may be used to identify individuals,  
CC for parentage testing, and in the genetic mapping of economic trait  
CC loci, or genes involved in the determination of economically important  
CC traits-esp. in cattle, to allow selective breeding.  
CC See also AA033501-34437.  
CC

SQ Sequence 35 BP; 3 A; 1 C; 15 G; 16 T; 0 other;

Query Match 0.5%; Score 19; DB 13; Length 35;  
Best Local Similarity 100.0%; Pred. No. 86;  
Matches 19; Conservative 0; Mismatches 0; Indels 0; Gaps 0;

QY 3844 tgctgctgtgtgtgtgt 3862  
|||  
DB 8 tgctgctgtgtgtgtgt 26

RESULT 19  
AA033669  
ID AA033669 standard; DNA; 37 BP.  
XX AA033669;  
AC  
XT 02-FEB-1993 (first entry)  
XX Microsatellite sequence from clone TGLA112.

```

Query Match          0.5%; Score 19; DB 13; Length 37;
Best Local Similarity 100.0%; Pred. NO. 86;
Matches 19; Conservative 0; Mismatches 0; Gaps 0;

OY      3857 gtcgtatgtcgtcgtcgtg 3875
          |||||
Db       19 gtcgtatgtcgtcgtcgtg 37

RESULT 20
AAT65709/c
AAT65709 standard; DNA; 46 BP.

XX AC AAT65709;
XX AC
DT DT
XX 17-JUN-1997 (first entry)
XX XX

DE Repeat sequence from polymorphic marker clone Mfd7.
XX KM
XX Polymorphism; repeat sequence; genetic marker; primer; amplification;
XX PCR; polymerase chain reaction; paternity; maternity; human; pedigree;
XX linkage analysis; genetic disease; animal; plant; breeding; locus;
XX hybridisation; chromosome; ds.
XX XX
OS Homo sapiens.
XX XX
XX USS582979-A.
XX XX
XX XX

```

	0.5%;	Score 19;	DB 18;	Length 46;
Query Match Similarity	100.0%;	Pred. No. 86;		
Matches 19; Conservative	0;	Mismatches	0;	Indels      0; Gaps    0
QY	3858	tgtgatatgcctgcctgcctc	3876	
DG	46	TGTGTATGTCGTGCTGTGT	28	

RESULT	21
AAQ34167	standard; DNA; 50 BP.
ID	AAQ34167
XX	
AC	AAQ34167;
XX	
DT	02-FEB-1993 (first entry)
XX	
DE	Sequence of a microsatellite from clone TGIAB5.
XX	
KM	PCR; selection; primers; OPTIPRIM; breeding; cattle; parentage
XX	
XX	genetic mapping; traits; amplification; ss.
OS	Bos taurus.
XX	
PN	WO9213102-A.
XX	
PD	06-AUG-1992.
XX	
PF	15-JAN-1992; 92WO-US00340.
XX	
PR	15-JAN-1991; 91US-0642342.
XX	
PA	(GENM-) GENMARK.
XX	
PI	Georges M, Maesey JM;
XX	
WR	WPI; 1992-284684/34.

XX Polymorphic bovine DNA markers - used in genetic identification,  
 PT gene mapping, and selective breeding  
 XX  
 PS Table 7; Page 397; 517pp; English.

CC The sequence is that of a bovine microsatellite sequence obt.  
 CC by screening a library of bovine MboI DNA fragments of between  
 CC 250 and 500 bp with an (AC)<sub>15</sub> and a (TC)<sub>15</sub> oligonucleotide probe.  
 CC One out of 50 clones cross-hybridised. Assuming independent  
 CC distribution of microsatellites and MboI sites, the frequency of  
 CC (16)<sup>n</sup> > 9 microsatellites in the bovine genome is estimated at >100,  
 CC 000. The sequence information for ca. 230 such bovine microsatellites  
 CC is summarised in the specification and indexed herein (see below).  
 CC The sequences upstream and downstream of the microsatellite sequence  
 CC were used to generate the required PCR primers for in vitro  
 CC amplification of the corresp. microsatellite (using the program  
 CC OPTIPRIM). The microsatellites may be used to identify individuals,  
 CC for parentage testing, and in the genetic mapping of economic trait  
 CC loci, or genes involved the determination of economically important  
 CC traits esp. in cattle, to allow selective breeding.  
 CC See also AAQ33501-34437.

XX Sequence 50 BP; 16 A; 0 C; 8 G; 26 T; 0 other;

SO

Query Match 0.5%; Score 19; DB 13; Length 50;  
 Best Local Similarity 100.0%; Pred. No. 85;  
 Matches 19; Conservative 0; Mismatches 0; Indels 0; Gaps 0;

OY 3852 tgcgtgtgtgtatgtgtgt 3870  
 |||||  
 Db 12 tgcgtgtgtgtatgtgtgt 30

RESULT 22  
 AAQ40668/C  
 ID AAQ40668 standard; CDNA; 24 BP.

XX AAQ40668;  
 XX

DT 06-AUG-1993 (first entry)

XX tPA-12 in vitro mutagenic oligomer.  
 DE  
 XX  
 XX Blood: tissue plasminogen activator; tPA; mutein; stability; tPA-12;  
 KM physiological; activity; pTB 1353; mutagenesis; plasmid; ss.  
 KW  
 XX  
 XX Synthetic.  
 OS  
 XX JP05076361-A.  
 PN  
 XX 30-MAR-1993.  
 PD  
 XX  
 XX 10-MAY-1991; 91JP-0105689.  
 PF  
 XX  
 XX 10-MAY-1990; 90JP-0118710.  
 PR  
 XX 25-DEC-1990; 90JP-0405848.  
 PS  
 XX (TAKE ) TAKEDA CHEM IND LTD.  
 PA  
 XX WPI; 1993-139567/17.  
 DR  
 XX  
 XX Tissue plasminogen activator mutein - useful for treating  
 PT myocardial infarction and cerebral thrombosis  
 PT  
 XX  
 XX Disclosure; Page 33; 92pp; Japanese.

XX This sequence is an oligomer which was used in an oligonucleotide  
 CC directed in vitro mutagenesis system for the production of tissue  
 CC plasminogen activator (tPA) mutein, tPA-12. The plasmid pTB 1353  
 CC was treated with this synthetic oligomer (see also AAQ40668). This  
 CC tPA mutein has good stability in blood and good physiological

CC activity.  
 XX  
 SO Sequence 24 BP; 6 A; 3 C; 9 G; 6 T; 0 other;

Query Match 0.4%; Score 18; DB 14; Length 24;  
 Best Local Similarity 100.0%; Pred. No. 2.6e+02;  
 Matches 18; Conservative 0; Mismatches 0; Indels 0; Gaps 0;

OY 616 tgcgaagcaccactact 633  
 |||||  
 Db 23 TTGCAAGCACCCTACTAT 6

RESULT 23  
 AA298502/C  
 ID AA298502 standard; DNA; 30 BP.

XX AA298502;  
 AC  
 XX

DT 19-JUN-2000 (first entry)

XX H. discus derived sequence #20.  
 DE  
 XX

KM Satellite sequence; DNA fragmentation; microsatellite DNA; DNA marker;  
 KM Hallotis discus; ss.  
 XX  
 XX Hallotis discus.  
 OS  
 XX WO200011156-A1.  
 PN  
 XX  
 XX 02-MAR-2000.  
 PD  
 XX  
 XX 01-JUL-1999; 99WO-JP03551.  
 PF  
 XX  
 XX 18-AUG-1998; 98JP-0232153.  
 PR  
 XX  
 XX (NORO ) JAPAN MIN AGRIC FORESTRY & FISHERIES.  
 PA  
 XX  
 XX Takahashi H, Sekino M;  
 PI  
 XX  
 XX WPI; 2000-224692/19.  
 DR  
 XX

PT Isolation of satellite sequences from genomic DNA for use as DNA  
 PT markers comprises isolating a library with high homogeneity by DNA  
 PT fragmentation -  
 PS  
 XX Example 5; Page 14; 35pp; Japanese.

CC The invention provides a novel method for isolation of satellite  
 CC sequences from genomic DNA that comprises fragmentation of the DNA by  
 CC a method which is not dependent on base sequences, then selection of  
 CC the satellite sequences from the obtained genomic library of high  
 CC homogeneity. The method is useful for the isolation of microsatellite  
 CC DNA sequences which can be used as DNA markers. The new method markedly  
 CC improves the efficiency of isolation of satellite sequences in  
 CC comparison to prior art methods which are reliant on base sequences.  
 CC Sequences AA298483-514 represent sequences from Hallotis discus, used in  
 CC the method of the invention.  
 CC  
 XX Sequence 30 BP; 15 A; 13 C; 0 G; 2 T; 0 other;

Query Match 0.4%; Score 18; DB 21; Length 30;  
 Best Local Similarity 100.0%; Pred. No. 2.6e+02;  
 Matches 18; Conservative 0; Mismatches 0; Indels 0; Gaps 0;

OY 3849 gtcgtgtgtgtatgt 3866  
 |||||  
 Db 19 GTCGTGTGTGTGTATGT 2

RESULT 24

AAQ34119  
ID AAQ34119 standard; DNA; 32 BP.  
XX  
AC AAQ34119;  
XX  
DT 02-FEB-1993 (first entry)  
XX  
DE Sequence of a microsatellite from clone TGLA67.  
XX  
XX PCR; selection; primers; OPTIPRIM; breeding; cattle; parentage;  
KM genetic mapping; traits; amplification; ss.  
XX  
OS Bos taurus.  
XX  
PN WO9213102-A.  
XX  
PD 06-AUG-1992.  
XX  
PF 15-JAN-1992; 92MO-US00340.  
XX  
PR 15-JAN-1991; 91US-0642342.  
XX  
XX (GENM-) GENMARK.  
XX  
PI Georges M, Massey JM;  
XX  
DR WPI; 1992-284684/34.  
XX  
PT Polymorphic bovine DNA markers - used in genetic identification,  
PT gene mapping, and selective breeding  
XX  
PS Table 7; Page 378; 517pp; English.  
XX  
CC The sequence is that of a bovine microsatellite sequence obtd.  
CC by screening a library of bovine MboI DNA fragments of between  
CC 250 and 500 bp with an (AC)<sub>15</sub> and a (TC)<sub>15</sub> oligonucleotide probe.  
CC One out of 50 clones cross-hybridised. Assuming independent  
CC distribution of microsatellites and MboI sites, the frequency of  
CC (T6)n > 9 microsatellites in the bovine genome is estimated at >100,  
CC 000. The sequence information for ca. 230 such bovine microsatellites  
CC is summarised in the specification and indexed herein (see below).  
CC The sequences upstream and downstream of the microsatellite sequence  
CC were used to generate the required PCR primers for in vitro  
CC amplification of the corresp. microsatellite (using the program  
CC OPTIPRIM). The microsatellites may be used to identify individuals,  
CC for parentage testing, and in the genetic mapping of economic trait  
CC loci, or genes involved the determination of economically important  
CC traits esp. in cattle, to allow selective breeding.  
CC See also AAQ33501-34437.  
XX  
XX Sequence 32 BP; 0 A; 1 C; 16 G; 15 T; 0 other;  
SQ

Query Match 0.4%; Score 18; DB 13; Length 32;  
Best Local Similarity 100.0%; Pred. No. 2.6e+02;  
Matches 18; Conservative 0; Mismatches 0; Indels 0; Gaps 0;

OY 3844 tctcgtgtgtgtgtgtg 3861.  
DB 15 tctcgtgtgtgtgtgtg 32

RESULT 25  
AAQ33698  
ID AAQ33698 standard; DNA; 37 BP.  
XX  
AC AAQ33698;  
XX  
DT 02-FEB-1993 (first entry)  
XX  
DE Microsatellite sequence from clone TGLA128.  
XX  
KM PCR; selection; primers; OPTIPRIM; breeding; cattle; parentage;

KM genetic mapping; traits; amplification; ss.  
XX  
OS Bos taurus.  
XX  
PN WO9213102-A.  
XX  
PD 06-AUG-1992.  
XX  
PF 15-JAN-1992; 92MO-US00340.  
XX  
PR 15-JAN-1991; 91US-0642342.  
XX  
XX (GENM-) GENMARK.  
XX  
PI Georges M, Massey JM;  
XX  
DR WPI; 1992-284684/34.  
XX  
PT Polymorphic bovine DNA markers - used in genetic identification,  
PT gene mapping, and selective breeding  
XX  
PS Table 7; Page 209; 517pp; English.  
XX  
CC The sequence is that of a bovine microsatellite sequence obtd. by  
CC screening a library of bovine MboI DNA fragments of between  
CC 250 and 500 bp with an (AC)<sub>15</sub> and a (TC)<sub>15</sub> oligonucleotide probe.  
CC One out of 50 clones cross-hybridised. Assuming independent  
CC distribution of microsatellites and MboI sites, the frequency of  
CC (T6)n > 9 microsatellites in the bovine genome is estimated at >100,  
CC 000. The sequence information for ca. 230 such bovine microsatellites  
CC is summarised in the specification and indexed herein (see below).  
CC The sequences upstream and downstream of the microsatellite sequence  
CC were used to generate the required PCR primers for in vitro  
CC amplification of the corresp. microsatellite (using the program  
CC OPTIPRIM). The microsatellites may be used to identify individuals,  
CC for parentage testing, and in the genetic mapping of economic trait  
CC loci, or genes involved the determination of economically important  
CC traits esp. in cattle, to allow selective breeding.  
CC See also AAQ33501-34437.  
XX  
XX Sequence 37 BP; 2 A; 1 C; 15 G; 19 T; 0 other;  
SQ

Query Match 0.4%; Score 18; DB 13; Length 37;  
Best Local Similarity 100.0%; Pred. No. 2.6e+02;  
Matches 18; Conservative 0; Mismatches 0; Indels 0; Gaps 0;

OY 3849 gtgtcgtgtgtgtgtatgt 3866  
DB 20 gtgtcgtgtgtgtgtatgt 37

RESULT 26  
AAT65788/C  
ID AAT65788 standard; DNA; 43 BP.  
XX  
AC AAT65788;  
XX  
DT 17-JUN-1997 (first entry)  
XX  
DE Repeat sequence from polymorphic marker clone Mcd117.  
XX  
KM Polymorphism; repeat sequence; genetic marker; primer; amplification;  
KM PCR; polymerase chain reaction; paternity; maternity; human; pedigree;  
KM linkage analysis; genetic disease; animal; plant; breeding; locus;  
KM hybridisation; chromosome; ds.  
XX  
OS Homo sapiens.  
XX  
PN US5582979-A.  
XX  
PD 10-DEC-1996.  
XX

PF 21-APR-1989; 89US-0341562.  
 XX  
 PR 05-SEP-1991; 91US-0754351.  
 PR 21-APR-1989; 89US-0341562.  
 PR 04-APR-1994; 94US-0222177.  
 XX  
 PA (MARS-) MARSHFIELD CLINIC.  
 XX  
 PI Weber JL.  
 XX  
 DR WPI; 1997-042299/04.  
 XX  
 PT Detection of polymorphic genetic markers of the form  
 (dc-da)n(dg-dt)n - using novel nucleic acid mols. as primers  
 PS  
 XX Claim 1; Column 13-14; 186pp; English.  
 CC The invention relates to the isolation of polymorphic repeat sequences  
 CC having the sequence (dc-da)n.(dg-dt)n which can be used as genetic  
 CC markers. Primers based on these sequences can be used to detect these  
 CC repeats, especially for use in e.g. paternity or maternity testing,  
 CC human genetic analysis such as linkage analysis of genetic disease,  
 CC commercial animal or plant breeding or pedigree analysis. Clones  
 CC containing the repeat sequences were isolated by hybridisation of  
 CC chromosome-specific phage libraries with a synthetic poly(dc-da) (dg-dt)  
 CC probe. Over 100 repeat blocks were isolated. The inserts from the  
 CC clones were amplified by primers AAT65798-T66047. Those clones where the  
 CC repeat sequence has been determined are shown in AAT65704-797. This  
 CC repeat sequence is from the marker clone Mdf117 which contains the  
 CC repeat sequence having the formula: GCAGCAACAT(AC)16.5.  
 XX  
 SQ Sequence 43 BP; 21 A; 19 C; 2 G; 1 T; 0 other:  
 XX  
 Query Match 0.4%; Score 18; DB 18; Length 43;  
 Best Local Similarity 100.0%; Pred. No. 2.5e+02;  
 Matches 18; Conservative 0; Mismatches 0; Indels 0; Gaps 0;  
 OY 3849 gtggtgtgtgtgtgtgtgt 3866  
 ||||||||||||||||||  
 Db 24 GTGCTGTGTGTGTGTGTGT 7  
 ||||||||||||||||||  
 RESULT 27  
 AA298507/C  
 ID AA298507 standard; DNA; 44 BP.  
 XX  
 AC AA298507;  
 XX  
 DT 19-JUN-2000 (first entry).  
 XX  
 DE H. discus derived sequence #25.  
 XX  
 KW Satellite sequence; DNA fragmentation; microsatellite DNA; DNA marker;  
 KW Halictis discus; ss.  
 XX  
 OS Halictis discus.  
 XX  
 PN NO200011156-A1.  
 XX  
 PD 02-MAR-2000.  
 XX  
 PF 01-JUL-1999; 99WO-JP03551.  
 XX  
 PR 18-AUG-1998; 98JP-0232153.  
 XX  
 PA (MORO) JAPAN MIN AGRIC FORESTRY & FISHERIES.  
 XX  
 PI Takahashi H, Sekino M;  
 XX  
 DR WPI; 2000-224692/19.  
 XX  
 PT Isolation of satellite sequences from genomic DNA for use as DNA

PT markers comprises isolating a library with high homogeneity by DNA  
 fragmentation -  
 XX  
 PS Example 5; Page 14; 35pp; Japanese.  
 XX  
 CC The invention provides a novel method for isolation of satellite  
 CC sequences from genomic DNA that comprises fragmentation of the DNA by  
 CC a method which is not dependent on base sequences, then selection of  
 CC the satellite sequences from the obtained genomic library of high  
 CC homogeneity. The method is useful for the isolation of microsatellite  
 CC DNA sequences which can be used as DNA markers. The new method markedly  
 CC improves the efficiency of isolation of satellite sequences in  
 CC comparison to prior art methods which are reliant on base sequences.  
 CC Sequences AA298483-514 represent sequences from Halictis discus, used in  
 CC the method of the invention.  
 XX  
 SQ Sequence 44 BP; 17 A; 20 C; 7 G; 0 U; 0 other:  
 XX  
 Query Match 0.4%; Score 18; DB 21; Length 44;  
 Best Local Similarity 100.0%; Pred. No. 2.5e+02;  
 Matches 18; Conservative 0; Mismatches 0; Indels 0; Gaps 0;  
 OY 3845 gtgctgtgtgtgtgtgtgt 3862  
 ||||||||||||||||||  
 Db 19 GTGCTGTGTGTGTGTGTGT 2  
 ||||||||||||||||||  
 RESULT 28  
 AAA80357  
 ID AAA80357 standard; DNA; 23 BP.  
 XX  
 AC AAA80357;  
 XX  
 DT 22-NOV-2000 (first entry)  
 XX  
 DE Human ASTH1 5' region polymorphic site, SEQ ID NO:103 (a).  
 XX  
 KW ASTH1 locus; ASTH1; human; chromosome 11p; asthma;  
 KW bronchial hyperactivity; ets family; transcription factor;  
 KW splice variant; genetic predisposition; polymorphism; antibody;  
 KW drug screening; prophylaxis; therapy; diagnosis;  
 KW single nucleotide polymorphism; SNP; ss.  
 XX  
 OS Homo sapiens.  
 XX  
 FH Key  
 FT Variation Location/Qualifiers  
 FT replace (12..13,TGTGTA)  
 FT /\*tag-a  
 XX  
 PN US6087485-A.  
 XX  
 PD 11-JUL-2000.  
 XX  
 PF 21-JAN-1998; 98US-0009913.  
 XX  
 PR 21-JAN-1997; 97US-0035663.  
 PR 01-JUL-1997; 97US-0051432.  
 XX  
 PA (AXYS-) AXYS PHARM INC.  
 XX  
 PI Galvin M, Miller A, North M, Cardon L, Buckler A;  
 PI Brooks-Wilson AR, Carey AH;  
 XX  
 DR WPI; 2000-505109/45.  
 XX  
 PT New nucleic acids other than naturally occurring chromosomes encoding  
 PT ASTH1 protein, for e.g. screening compositions that modulate expression  
 PT or function of ASTH1 proteins or as diagnostics for genetic  
 PT predisposition to asthma  
 XX  
 PS Examples: Column 41-42; 131pp; English.







Matches 17; Conservative 0; Mismatches 0; Indels 0; Gaps 0;

OY 3849 gtgtgtgtgtgtgtgtgt 3865  
 ||||||||||||||||  
 DB 17 GTGTGTGTGTGTGTATG 1

## RESULT 33

AA268141/C  
 ID AA268141 standard; DNA: 47 BP.

AC AA268141;  
 XX

DT 10-SEP-2001 (first entry)  
 XX

DE Human map-related biallelic marker SEQ ID NO:2488.  
 XX

KM Human genome; biallelic marker; high density disequilibrium map;  
 KW genomic map; haplotype; phenotype; polymorphic base; genotyping;  
 KM haplotyping; hybridisation; identification; characterisation;  
 KM diagnosis; single nucleotide polymorphism; SNP; ds.

XX Homo sapiens.  
 OS

FT Key Location/Qualifiers  
 FT variation replace(24,C)  
 FT /\*tag- a  
 FT /standard\_name= "single nucleotide polymorphism"

PN WO954500-A2.  
 XX

PD 28-OCT-1999.  
 XX

PF 21-APR-1999; 99MO-IB00822.  
 XX

PR 21-APR-1998; 98US-0082614.  
 PR 23-NOV-1998; 98US-0109732.  
 XX

PA (GEST ) GENSET.  
 XX

PI Cohen D, Blumenfeld M, Chumakov I;  
 XX

DR WPI; 2000-013267/01.  
 XX

PT Novel biallelic markers used to construct a high density disequilibrium  
 map of the human genome.  
 XX

PS Claim 3; Page 761; 2745pp; English.  
 XX

CC AA265654 to AA269578 represent human biallelic markers from the present  
 CC invention, which contain a polymorphic base at position 24 of their  
 CC nucleotide sequences. AA269579 to AA277440 represent amplification  
 CC primers for the biallelic markers. The biallelic markers of the  
 CC invention have a variety of uses: they can be used for high density  
 CC mapping of the human genome, and in complex association studies and  
 CC haplotyping studies which are useful in determining the genetic basis  
 CC for disease states. Compositions and methods of the invention can also  
 CC be useful for the identification of the targets for the development of  
 CC pharmaceutical agents and diagnostic methods, as well as the  
 CC characterisation of the differential efficacious responses to and side  
 CC effects from pharmaceutical agents acting on a disease as well as other  
 CC treatment.  
 CC N.B. The SEQ ID NOS 2852, 2913, 2974, 3035, 3096, 3157, 3227, 3297  
 CC and 3367, are not actually given a sequence in the Sequence Listing  
 CC from the present invention.  
 XX

SO Sequence 47 BP; 23 A; 9 C; 2 G; 13 T; 0 other;

Query Match 0.4%; Score 17; DB 21; Length 47;  
 Best Local Similarity 100.0%; Pred. No. 7.6e+02;  
 Matches 17; Conservative 0; Mismatches 0; Indels 0; Gaps 0;

OY 3862 tatgtgtgtgtgtat 3878  
 ||||||||||||||||  
 DB 25 TATGTGTGTGTGTATG 9

## RESULT 34

AA034122  
 ID AA034122 standard; DNA: 49 BP.

AC AA034122;  
 XX

DT 02-FEB-1993 (first entry)  
 XX

DE Sequence of a microsatellite from clone TGIAB6.  
 XX

KM PCR; selection; primers; OPTIPRIM; breeding; cattle; parentage;  
 KW genetic mapping; traits; amplification; ss.  
 XX

OS Bos taurus.  
 XX

PN WO9213102-A.  
 XX

PD 06-AUG-1992.  
 XX

PF 15-JAN-1992; 92MO-US00340.  
 XX

PR 15-JAN-1991; 91US-0642342.  
 XX

PA (GENM-) GENMARK.  
 XX

PI Georges M, Massey JM;  
 XX

DR WPI; 1992-284684/34.  
 XX

PT Polymorphic bovine DNA markers - used in genetic identification,  
 gene mapping, and selective breeding  
 XX

PS Table 7; Page 379; 517pp; English.  
 XX

CC The sequence is that of a bovine microsatellite sequence obtd.  
 CC by screening a library of bovine MboI DNA fragments of between  
 CC 250 and 500 bp with an (AC)<sub>15</sub> and a (TC)<sub>15</sub> oligonucleotide probe.  
 CC One out of 50 clones cross-hybridised. Assuming independent  
 CC distribution of microsatellites and MboI sites, the frequency of  
 CC (76)n >9 microsatellites in the bovine genome is estimated at >100,  
 CC 000. The sequence information for ca. 230 such bovine microsatellites  
 CC is summarised in the specification and indexed herein (see below).  
 CC The sequences upstream and downstream of the microsatellite sequence  
 CC were used to generate the required PCR primers for in vitro  
 CC amplification of the corresp. microsatellite (using the program  
 CC OPTIPRIM). The microsatellites may be used to identify individuals,  
 CC for parentage testing, and in the genetic mapping of economic trait  
 CC loci, or genes involved in the determination of economically important  
 CC traits esp. in cattle, to allow selective breeding.  
 CC See also AA033501-34437.  
 XX

SO Sequence 49 BP; 7 A; 0 C; 17 G; 25 T; 0 other;

Query Match 0.4%; Score 17; DB 13; Length 49;  
 Best Local Similarity 100.0%; Pred. No. 7.6e+02;  
 Matches 17; Conservative 0; Mismatches 0; Indels 0; Gaps 0;

OY 3860 tgatgtgtgtgtgtgt 3876  
 ||||||||||||||||  
 DB 1 tgatgtgtgtgtgtgtgt 17

RESULT 35  
 AAT30412  
 ID AAT30412 standard; DNA: 19 BP.  
 AC AAT30412;

```

XX 28-JAN-1997 (first entry)
XX Compound simple sequence repeat primer (GT)7.5(AT)2.
XX
XX Detection: polymorphism; perfect compound simple sequence repeat;
XX adaptor directed primer; genome: genetic; fingerprinting;
XX amplified fragment length polymorphism assay;
XX microsatellite region; genetic trait marking;
XX germplasm comparisons; compound; ss.
XX Synthetic.
XX
XX WO9617082-A2.
XX
XX 06-JUN-1996.
XX
XX 21-NOV-1995; 95WO-0515150.
XX
XX 28-NOV-1994; 94US-0346456.
XX
XX (DUPO) DU PONT DE NEMOURS & CO E. I.
XX
XX Morgante M, Vogel JM;
XX
XX WPI: 1996-277795/28.
XX
XX Modified amplified fragment length polymorphism assay - for
XX detection of polymorphism esp. in microsatellite regions
XX
XX Example 2; Page 84; 173pp; English.
XX
XX Detecting polymorphisms between 2 nucleic acid samples, esp. in
XX microsatellite regions, comprises digesting the nucleic acid to
XX generate fragments, ligating adaptor segments to their ends,
XX amplifying them using primer directed amplification and comparing
XX the prods. to detect differences. The primers used in the
XX amplification comprise a primer consisting of a perfect cpd. simple
XX sequence repeat (SSR), and an adaptor directed primer, comprising a
XX sequence complementary to an adaptor segment. The present sequence
XX is an example of a compound SSR primer.
XX The method represents a modified amplified fragment length
XX polymorphism assay, which is partic. useful for genome
XX fingerprinting, i.e. for genetic trait marking and germplasm
XX comparisons.
XX
XX Sequence 19 BP; 2 A; 0 C; 7 G; 10 T; 0 other;
XX
XX
XX Query Match 0.4%; Score 16; DB 17; Length 19;
XX Best Local Similarity 100.0%; Pred. No. 2.3e+03;
XX Matches 16; Conservative 0; Mismatches 0; Indels 0; Gaps 0;
XX
XX QY 3849 gtgtgtgtgtgtgtat 3864
XX |||||||||||||||
XX DB 2 gtgtgtgtgtgtgtat 17
XX
XX
XX RESULT 36
XX AA041067/c
XX ID AA041067 standard; DNA; 19 BP.
XX
XX AA041067;
XX
XX 25-SEP-1998 (first entry)
XX
XX Primer TEL:114019 for abnormality detection.
XX
XX PCR primer: chromosomal abnormality; abnormality detection; leukemia;
XX lymphoma; carcinoma; adenocarcinoma; sarcoma; glioma; neuroblastoma;
XX medullablastoma; malignant melanoma; malignant neoplastic condition; ss.
XX
XX Synthetic.
XX
XX OS

```

```

OS Homo sapiens.
XX
XX WO9824928-A2.
XX
XX 11-JUN-1998.
XX
XX 08-DEC-1997; 97WO-DK00556.
XX
XX 06-DEC-1996; 96DK-0001401.
XX
XX (PALL/) PALLISGAARD N.
XX
XX Hokland P, Pallisgaard N;
XX
XX WPI: 1998-333344/29.
XX
XX Detection of chromosomal abnormalities - by subjecting patient
XX sample nucleic acids to a multiplex molecular amplification
XX procedure using primers specific for characteristic nucleic acid
XX sequence
XX
XX Claim 73; Page 107; 126pp; English.
XX
XX This sequence represents a primer used in the method of the invention for
XX the detection of the presence or absence of chromosomal abnormalities,
XX each abnormality being associated with a condition in a subject and each
XX being defined by at least one characteristic nucleic acid sequence. The
XX method comprises: (a) obtaining a sample of nucleic acids derived from a
XX subject which may harbour one of the chromosomal abnormalities;
XX (b) subjecting the sample to a multiplex molecular amplification (MMA)
XX procedure, where a number of the characteristic sequences, if present in
XX a sufficient amount, will be amplified; (c) retrieving the product(s)
XX from step (b), and detecting the presence and/or absence of an amplicon
XX characteristic of the abnormal sequences to detect the presence or
XX absence of corresponding chromosomal abnormalities; where the MMA
XX procedure comprises the use of at least 7 mutually distinct primers (MDP)
XX in one single reaction mixture, each of the primers defining an end of at
XX least one characteristic nucleic acid sequence, and where at least one of
XX the primers defines the first end of at least two characteristic nucleic
XX acid sequences; the characteristic nucleic acid sequences each being
XX determined in their opposite ends by MDP selected from the remainder of
XX the MDP. The methods can be used for detecting chromosomal abnormalities
XX associated with diseases including numerous leukaemia's, lymphoma's,
XX carcinoma's, adenocarcinoma's, sarcoma's, glioma's, neuroblastoma's,
XX medullablastoma, malignant melanoma, and malignant neoplastic conditions.
XX
XX Sequence 19 BP; 4 A; 7 C; 3 G; 5 T; 0 other;
XX
XX
XX Query Match 0.4%; Score 16; DB 19; Length 19;
XX Best Local Similarity 100.0%; Pred. No. 2.3e+03;
XX Matches 16; Conservative 0; Mismatches 0; Indels 0; Gaps 0;
XX
XX QY 1701 tggacatgaagtcgc 1716
XX |||||||||||||||
XX DB 19 TGGACATGAAGTCGC 4
XX
XX
XX RESULT 37
XX AA048546/c
XX ID AA048546 standard; DNA; 20 BP.
XX
XX AA048546;
XX
XX 22-FEB-1994 (first entry)
XX
XX HPV E6/7 region probe.
XX
XX Human papilloma virus; HPV; E6; E7; benign; malignant; probe; ss.
XX
XX Synthetic.
XX
XX JP05192200-A.
XX
XX PN

```

XX 03-AUG-1993.  
PD  
XX 19-AUG-1991; 91JP-0230839.  
PF  
XX 20-AUG-1990; 90JP-0217067.  
PR  
XX (TAKI ) TAKARA SHUZO CO LTD.  
PA  
XX WPI; 1993-277497/35.  
DR  
XX  
XX  
PT Detecting benign and/or malignant human papilloma virus - by  
PT detecting DNA sequence of E6 and/or E7 region of human papilloma  
virus  
XX  
XX  
PS Disclosure; Page 14; 18pp; Japanese.  
XX  
XX  
CC The probe is used to detect benign and/or malignant human papilloma  
CC virus. The probe binds to the E6 and/or E7 region of the virus.  
XX  
SQ Sequence 20 BP; 6 A; 6 C; 3 G; 5 T; 0 other;

Query Match 0.4%; Score 16; DB 14; Length 20;  
Best Local Similarity 100.0%; Pred. No. 2.3e+03;  
Matches 16; Conservative 0; Mismatches 0; Indels 0; Gaps 0;  
QY 2382 tgcacagcagtgtaag 2397  
DB 16 TGTCACAGAGTGAAG.1

RESULT 38  
AAT30427/c  
ID AAT30427 standard; DNA: 20 BP.  
XX  
XX AAT30427;  
AC  
XX 28-JAN-1997 (first entry)  
XX  
XX Compound simple sequence repeat primer (CA)4.5(TA)7.5.  
DE  
XX  
XX  
XX Detection: polymorphism; perfect compound simple sequence repeat;  
KM adaptor directed primer; genome; genetic; fingerprinting;  
KM amplified fragment length polymorphism assay;  
KM microsatellite region; genetic trait marking;  
KM germlasm comparisons; compound; ss.  
XX  
OS Synthetic.  
XX  
XX W09617082-A2.  
PN  
XX  
XX 06-JUN-1996.  
PD  
XX  
XX 21-NOV-1995; 95WO-US15150.  
PF  
XX 28-NOV-1994; 94US-0346456.  
PR  
XX  
XX (DUPO ) DU PONT DE NEMOURS & CO E. I.  
PA  
XX  
XX Morgante M, Vogel JM;  
PI  
XX  
XX WPI; 1996-277795/28.  
DR  
XX  
XX  
XX Modified amplified fragment length polymorphism assay - for  
PT detection of polymorphism esp. in microsatellite regions  
XX  
XX  
PS Disclosure; Fig 1c; 173pp; English.  
XX  
XX  
CC Detecting polymorphisms between 2 nucleic acid samples, esp. in  
CC microsatellite regions, comprises digesting the nucleic acid to  
CC generate fragments, ligating adaptor segments to their ends,  
CC amplifying them using primer directed amplification and comparing

CC the probe to detect differences. The primers used in the  
CC amplification comprise a primer consisting of a perfect cpd, simple  
CC sequence repeat (SSR), and an adaptor directed primer, comprising a  
CC sequence complementary to an adaptor segment. The present sequence  
CC is an example of a compound SSR primer.  
CC The method represents a modified amplified fragment length  
CC polymorphism assay, which is partic. useful for genome  
CC fingerprinting, i.e. for genetic trait marking and germlasm  
CC comparisons.  
XX  
SQ Sequence 20 BP; 10 A; 7 C; 0 G; 3 T; 0 other;

Query Match 0.4%; Score 16; DB 17; Length 20;  
Best Local Similarity 100.0%; Pred. No. 2.3e+03;  
Matches 16; Conservative 0; Mismatches 0; Indels 0; Gaps 0;  
QY 3849 gtgtgtgtgtgtgtat 3864  
DB 19 GTGTGTGTGTGTGTAT 4

RESULT 39  
AAS09069  
ID AAS09069 standard; DNA: 20 BP.  
XX  
XX AAS09069;  
AC  
XX 26-SEP-2001 (first entry)  
XX  
XX  
XX Human MEK2 antisense oligonucleotide 113875.  
DE  
XX  
XX Human; mitogen-activated protein kinase-kinase kinase 2; MAP; MEK2;  
XX MEK kinase 2; MAP/ERK kinase kinase 2; immunological disorder;  
XX inflammatory disorder; hyperproliferative disorder; cancer; antisense;  
XX phosphorothioate; ss.  
XX  
XX Homo sapiens.  
OS  
XX  
XX  
XX Key Location/Qualifiers  
FH modified\_base 1..20  
FT /tag- a  
FT /mod\_base- "OTHER"  
FT /note- "OTHER- phosphorothioate internucleotide linkages.  
FT Some bases especially bases 1-5 and bases 16-20  
FT are 2'-methoxyethyl (2'-MOE) bases, bases 6-15  
FT are 2'-deoxynucleotides and all cytidine bases  
FT are 5'-methylcytidines"  
XX  
XX  
XX W0200152863-A1.  
PN  
XX  
XX 26-JUL-2001.  
PD  
XX  
XX 16-JAN-2001; 2001WO-US01361.  
PF  
XX 20-JAN-2000; 2000US-0486744.  
PR  
XX  
XX (ISIS-) ISIS PHARM INC.  
PA  
XX  
XX Monia BP, Gaarde WA, Ward DT, Freiler SM, Wyatt JR;  
PI  
XX  
XX WPI; 2001-442246/47.  
DR  
XX  
XX  
XX Antisense compound 8 to 30 nucleobases in length targeted to a nucleic  
PT acid molecule encoding MEK2, useful for the treatment of an  
PT immunological, inflammatory or hyperproliferative disorder -  
XX  
XX  
PS Claim 3; Page 79; 105pp; English.  
XX  
XX  
XX The present sequence for human MEK2 antisense oligonucleotide 113875  
CC is 1 of various novel human mitogen-activated protein (MAP)  
CC kinase kinase kinase 2 (MEK2, also known as MEK kinase 2 and  
CC MAP/ERK kinase kinase 2) antisense oligonucleotides (AAS09045-AAS09122)

CC which specifically hybridise with and inhibit the expression of MEKK2.  
 CC The antisense oligonucleotides can be used in a composition to modulate  
 CC the expression of MEKK2 (AAU03598). The antisense oligonucleotides are  
 CC useful for inhibiting the expression of MEKK2 in the treatment of  
 CC immunological disorders, inflammatory disorders and hyperproliferative  
 CC disorders e.g. cancer.  
 XX

SQ Sequence 20 BP; 2 A; 5 C; 4 G; 9 T; 0 other;

Query Match 0.4%; Score 16; DB 22; Length 20;  
 Best Local Similarity 100.0%; Pred. No. 2.3e+03;  
 Matches 16; Conservative 0; Mismatches 0; Indels 0; Gaps 0;

OY 2545 tctggttctctgcaag 2560  
 |||||  
 DB 5 tctggttctctgcaag 20

RESULT 40

AA03579/c  
 ID AAC83579 standard; DNA; 23 BP.

XX AAC83579;

DT 28-FEB-2001 (first entry)

DE Human FMR1 gene triplet repeat PCR primer NM-B5-for.

XX Human; FMR1; Fragile X syndrome; methylation; diagnosis;

KW chromosome Xq27.3; PCR primer; ss.

XX Homo sapiens.

XX US6143504-A.

XX 07-NOV-2000.

PF 27-OCT-1999; 99US-0429499.

XX 27-OCT-1999; 99US-0429499.

PA (ARCH-) ARCH DEV CORP.

PI Das S, Ledbetter DH;

XX WPI; 2001-006432/01.

PT Determining methylation state of FMR1 gene promoter for diagnosing  
 PT fragile X syndrome in males involves denaturing DNA sample, subjecting  
 PT DNA to bisulfite modification, amplifying DNA and detecting products -

XX claim 17; Column 31; 20pp; English.

CC The present invention describes a novel method of diagnosing Fragile X  
 CC syndrome using a PCR-based method of methylation analysis. The FMR1 gene  
 CC promoter, located at chromosome Xq27.3, is composed of a CGG  
 CC trinucleotide repeat. The expansion of this repeat leads to a premutation  
 CC and then a full mutation, the latter of which is likely to cause the  
 CC methylation of a nearby CpG island, causing the fragile X syndrome  
 CC phenotype. This method is useful in the design of appropriate therapies  
 CC and counselling for affected individuals and carriers.  
 XX

SQ Sequence 23 BP; 11 A; 10 C; 0 G; 2 T; 0 other;

Query Match 0.4%; Score 16; DB 22; Length 23;  
 Best Local Similarity 100.0%; Pred. No. 2.3e+03;  
 Matches 16; Conservative 0; Mismatches 0; Indels 0; Gaps 0;

OY 3855 gctgtgtatgtgtgt 3870  
 |||||  
 DB 16 gctgtgtatgtgtgt 1

RESULT 41

AA298498/c  
 ID AA298498 standard; DNA; 24 BP.

XX AA298498;

DT 19-JUN-2000 (first entry)

DE H. discus derived sequence #16.

XX Satellite sequence; DNA fragmentation; microsatellite DNA; DNA marker.

KW Hallois discus; ss.

XX Hallois discus.

XX WO200011156-A1.

XX 02-MAR-2000.

PF 01-JUL-1999; 99WO-JP03551.

XX 18-AUG-1998; 98JP-0232153.

PA (NORO) JAPAN MIN AGRIC FORESTRY & FISHERIES.

PI Takahashi H, Sekino M;

XX WPI; 2000-224692/19.

PT Isolation of satellite sequences from genomic DNA for use as DNA  
 PT markers comprises isolating a library with high homogeneity by DNA  
 PT fragmentation -

XX Example 5; Page 14; 35pp; Japanese.

CC The invention provides a novel method for isolation of satellite  
 CC sequences from genomic DNA that comprises fragmentation of the DNA by  
 CC a method which is not dependent on base sequences, then selection of  
 CC the satellite sequences from the obtained genomic library of high  
 CC homogeneity. The method is useful for the isolation of microsatellite  
 CC DNA sequences which can be used as DNA markers. The new method markedly  
 CC improves the efficiency of isolation of satellite sequences in  
 CC comparison to prior art methods which are reliant on base sequences.  
 CC Sequences AA298483-514 represent sequences from Hallois discus, used in  
 CC the method of the invention.  
 XX

SQ Sequence 24 BP; 8 A; 12 C; 4 G; 0 U; 0 other;

Query Match 0.4%; Score 16; DB 21; Length 24;  
 Best Local Similarity 100.0%; Pred. No. 2.3e+03;  
 Matches 16; Conservative 0; Mismatches 0; Indels 0; Gaps 0;

OY 3847 gctgtgtgtgtgtgt 3862  
 |||||  
 DB 19 GCGTGTGTGTGTGTGT 4

RESULT 42

AA034131  
 ID AA034131 standard; DNA; 26 BP.

XX AA034131;

DT 02-FEB-1993 (first entry)

DE Sequence of a microsatellite from clone TGA170B.

XX PCR; selection; primers; OPTIPRIM; breeding; cattle; parentage;  
 XX genetic mapping; traits; amplification; ss.  
 XX

OS Bos taurus.  
 XX  
 PN MO9213102-A.  
 XX  
 PD 06-AUG-1992.  
 XX  
 PF 15-JAN-1992; 92MO-US00340.  
 XX  
 PR 15-JAN-1991; 91US-0642342.  
 XX  
 PA (GENM-) GENMARK.  
 XX  
 PI Georges M, Massey JM;  
 XX  
 DR WPI; 1992-284684/34.  
 XX  
 PT Polymorphic bovine DNA markers - used in genetic identification,  
 XX gene mapping, and selective breeding  
 XX  
 PS Table 7; Page 383; 517pp; English.  
 XX  
 CC The sequence is that of a bovine microsatellite sequence obtd.  
 CC by screening a library of bovine MbOI DNA fragments of between  
 CC 250 and 500 bp with an (AC)<sub>15</sub> and a (TC)<sub>15</sub> oligonucleotide probe.  
 CC One out of 50 clones cross-hybridised. Assuming independent  
 CC distribution of microsatellites and MbOI sites, the frequency of  
 CC (76)n > 9 microsatellites in the bovine genome is estimated at >100,  
 CC 000. The sequence information for ca. 230 such bovine microsatellites  
 CC is summarised in the specification and indexed herein (see below).  
 CC The sequences upstream and downstream of the microsatellite sequence  
 CC were used to generate the required PCR primers for in vitro  
 CC amplification of the corresp. microsatellite (using the program  
 CC OPTIPRIM). The microsatellites may be used to identify individuals,  
 CC for parentage testing, and in the genetic mapping of economic trait  
 CC loci, or genes involved in the determination of economically important  
 CC traits esp. in cattle, to allow selective breeding.  
 CC See also AAQ33501-34437.  
 CC  
 XX  
 SQ Sequence 26 BP; 2 A; 1 C; 12 G; 11 T; 0 other;

Query Match 0.4%; Score 16; DB 13; Length 26;  
 Best Local Similarity 100.0%; Pred. No. 2.3e+03;  
 Matches 16; Conservative 0; Mismatches 0; Indels 0; Gaps 0;

OY 3847 gcgtgtgtgtgtgtgt 3862  
 |||  
 DB 7 gcgtgtgtgtgtgtgt 22

RESULT 43  
 AAQ33740  
 ID AAQ33740 standard; DNA; 27 BP.  
 XX  
 AC AAQ33740;  
 XX  
 DT 02-FEB-1993 (first entry)  
 XX  
 DE Microsatellite sequence from clone TGLA154.  
 XX  
 KW PCR; selection; primers; OPTIPRIM; breeding; cattle; parentage;  
 KW genetic mapping; traits; amplification; ss.  
 XX  
 OS Bos taurus.  
 XX  
 PN WO9213102-A.  
 XX  
 PD 06-AUG-1992.  
 XX  
 PF 15-JAN-1992; 92MO-US00340.  
 XX  
 PR 15-JAN-1991; 91US-0642342.  
 XX

PA (GENM-) GENMARK.  
 XX  
 PI Georges M, Massey JM;  
 XX  
 DR WPI; 1992-284684/34.  
 XX  
 PT Polymorphic bovine DNA markers - used in genetic identification,  
 XX gene mapping, and selective breeding  
 XX  
 PS Table 7; Page 226; 517pp; English.  
 XX  
 CC The sequence is that of a bovine microsatellite sequence obtd. by  
 CC screening a library of bovine MbOI DNA fragments of between  
 CC 250 and 500 bp with an (AC)<sub>15</sub> and a (TC)<sub>15</sub> oligonucleotide probe.  
 CC One out of 50 clones cross-hybridised. Assuming independent  
 CC distribution of microsatellites and MbOI sites, the frequency of  
 CC (76)n > 9 microsatellites in the bovine genome is estimated at >100,  
 CC 000. The sequence information for ca. 230 such bovine microsatellites  
 CC is summarised in the specification and indexed herein (see below).  
 CC The sequences upstream and downstream of the microsatellite sequence  
 CC were used to generate the required PCR primers for in vitro  
 CC amplification of the corresp. microsatellite (using the program  
 CC OPTIPRIM). The microsatellites may be used to identify individuals,  
 CC for parentage testing, and in the genetic mapping of economic trait  
 CC loci, or genes involved in the determination of economically important  
 CC traits esp. in cattle, to allow selective breeding.  
 CC See also AAQ33501-34437.  
 CC  
 XX  
 SQ Sequence 27 BP; 2 A; 0 C; 12 G; 13 T; 0 other;

Query Match 0.4%; Score 16; DB 13; Length 27;  
 Best Local Similarity 100.0%; Pred. No. 2.3e+03;  
 Matches 16; Conservative 0; Mismatches 0; Indels 0; Gaps 0;

OY 3849 gtgtgtgtgtgtgtgt 3864  
 |||  
 DB 4 gtgtgtgtgtgtgtgt 19

RESULT 44  
 AA161970/C  
 ID AA161970 standard; DNA; 27 BP.  
 XX  
 AC AA161970;  
 XX  
 DT 16-OCT-2001 (first entry)  
 XX  
 DE Soybean 240017 region G3 DNA forward primer, SEQ ID NO: 601.  
 XX  
 KW Soybean; antihelminthic; gene therapy; soybean cyst nematode; SCN;  
 KW SCN resistance; ring1; Ring4; SCN resistant allele; plant breeding;  
 KW 240017 region G3; 318013 region A3; 515002 region G2; PCR primer; ss.  
 XX  
 OS Glycine max.  
 XX  
 PN WO200151627-A2.  
 XX  
 PD 19-JUL-2001.  
 XX  
 PF 05-JAN-2001; 2001WO-US00552.  
 XX  
 PR 07-JAN-2000; 2000US-0174880.  
 XX  
 PA (MONS ) MONSANTO CO.  
 XX  
 PI Hauge BM, Wang ML, Parsons JD, Parnell LD;  
 XX  
 DR WPI; 2001-425872/45.  
 XX  
 PT New purified nucleic acid for producing a soybean plant having soybean  
 PT cyst nematode resistance and for use in plant breeding programs -  
 XX

PS Claim 25; Page 1178; 1353bp; English.  
 XX  
 CC The invention relates to nucleic acid molecules from regions of the  
 CC soybean genome which are associated with soybean cyst nematode (SCN)  
 CC resistance. The nucleic acids are used to transform plants, and can  
 CC produce soybean plants having an rhg1 or an Rhg4 SCN resistant allele.  
 CC The nucleic acids can be used for investigating rhg1 or Rhg4 haplotypes  
 CC of soybean plants and for introgressing SCN resistance or partial SCN  
 CC resistance into soybean plants. They can also be used in plant breeding  
 CC programmes. The invention also relates to proteins encoded by such  
 CC nucleic acid molecules, as well as antibodies capable of recognising a  
 CC these proteins. The present sequence is a primer used to amplify a  
 CC region of the soybean genome.  
 SQ Sequence 27 BP; 12 A; 11 C; 0 G; 4 T; 0 other;

Query Match 0.4%; Score 16; DB 22; Length 27;  
 Best Local Similarity 100.0%; Pred. No. 2.3e+03;  
 Matches 16; Conservative 0; Mismatches 0; Indels 0; Gaps 0;  
 QY 3849 gtcgtgtgtgtgtgtat 3864  
 DB 22 gtcgtgtgtgtgtgtat 7

RESULT 45  
 AAQ33761  
 ID AAQ33761 standard; DNA; 34 BP.  
 XX  
 AC AAQ33761;  
 XX  
 DT 02-FEB-1993 (first entry)  
 XX  
 DE Microsatellite sequence from clone TGLA170.  
 XX  
 KW PCR; selection; primers; OPTIPRM; breeding; cattle; parentage;  
 KW genetic mapping; traits; amplification; ss.  
 XX  
 OS Bos taurus.  
 XX  
 PN W09213102-A.  
 XX  
 PD 06-AUG-1992.  
 XX  
 PF 15-JAN-1992; 92WC-US00340.  
 XX  
 PR 15-JAN-1991; 91US-0642342.  
 XX  
 PA (GENM-) GENMARK.  
 XX  
 PI Georges M, Massey JM;  
 XX  
 DR WPI; 1992-284684/34.  
 XX  
 PT Polymorphic bovine DNA markers - used in genetic identification,  
 PT gene mapping, and selective breeding  
 XX  
 PS Table 7; Page 234; 517bp; English.  
 XX  
 CC The sequence is that of a bovine microsatellite sequence obt'd. by  
 CC screening a library of bovine MboI DNA fragments of between  
 CC 250 and 500 bp with an (AC)<sub>15</sub> and a (TC)<sub>15</sub> oligonucleotide probe.  
 CC One out of 50 clones cross-hybridised. Assuming independent  
 CC distribution of microsatellites and MboI sites, the frequency of  
 CC (T6)<sub>n</sub> >9 microsatellites in the bovine genome is estimated at >100,  
 CC 000. The sequence information for ca. 230 such bovine microsatellites  
 CC is summarised in the specification and indexed herein (see below).  
 CC The sequences upstream and downstream of the microsatellite sequence  
 CC were used to generate the required PCR primers for in vitro  
 CC amplification of the corresp. microsatellite (using the program  
 CC OPTIPRM). The microsatellites may be used to identify individuals,  
 CC for parentage testing, and in the genetic mapping of economic trait

CC loci, or genes involved the determination of economically important  
 CC traits esp. in cattle, to allow selective breeding.  
 CC See also AAQ33501-34437.  
 XX  
 SQ Sequence 34 BP; 0 A; 2 C; 17 G; 15 T; 0 other;

Query Match 0.4%; Score 16; DB 13; Length 34;  
 Best Local Similarity 100.0%; Pred. No. 2.3e+03;  
 Matches 16; Conservative 0; Mismatches 0; Indels 0; Gaps 0;  
 QY 3847 gcgtgtgtgtgtgtgt 3862  
 DB 13 gcgtgtgtgtgtgtgt 28

Search completed: April 19, 2002, 22:03:19  
 Job time: 9828 sec

---